



Electronic Pneumatic Injection-Assisted Dermal Drug Delivery Visualized by Ex Vivo Confocal Microscopy

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Background and Objectives: Electronic pneumatic injection (EPI) is a technique for dermal drug delivery, which is increasingly being used in clinical practice. However, only few studies have been reported on cutaneous drug distribution and related clinical endpoints. We aimed to visualize the immediate cutaneous drug distribution, changes in skin architecture, and related clinical endpoint of EPI.

Study Design/Materials and Methods: Acridine orange (AO) solution was administered to *ex vivo* porcine skin by EPI at pressure levels from 4 to 6 bar with a fixed injection volume of 50 μ l and nozzle size of 200 μ m. Immediate cutaneous distribution was visualized using *ex vivo* confocal microscopy (EVCN). Changes in skin architecture were visualized using both EVCN and hematoxylin and eosin-stained cryosections.

Results: The defined immediate endpoint was a clinically visible papule formation on the skin. The pressure threshold to consistently induce a papule was 4 bar, achieving delivery of AO to the deep dermis (2319 μ m axial and 5944 μ m lateral distribution). Increasing the pressure level to 6 bar did not lead to significant differences in axial and lateral dispersion ($P=0.842$, $P=0.905$; respectively). A distinctively hemispherical distribution pattern was identified. Disruption of skin architecture occurred independently of pressure level, and consisted of sub-epidermal clefts, dermal vacuoles, and fragmented collagen.

Conclusions: This is the first study to relate a reproducible clinical endpoint to EPI-assisted immediate drug delivery using EVCN. An EPI-induced skin papule indicates dermal drug delivery throughout all layers of the dermis, independent of pressure level settings. *Lasers Surg. Med.* © 2020 The Authors. *Lasers in Surgery and Medicine* published by Wiley Periodicals LLC

Key words: dermatology; skin; pneumatic device; needle-free injection; electronically-controlled; drug delivery; *ex vivo* confocal microscopy; device; biodistribution

INTRODUCTION

The skin functions as an effective barrier against external influences. When intending to deliver drugs

intradermally, conventional syringe-needle injection is the standard of care technique to overcome this barrier. Despite their low cost, wide availability, and ease of use, needle injections have several disadvantages including pain, needle-phobia (24% of adults [1]), and the risk of needle stick injuries. To overcome these restraints, other physical delivery techniques have been developed to provide effective intradermal drug delivery [2]. These techniques include laser-assisted drug delivery, micro-needling, electrochemotherapy, iontophoresis, and jet injectors [3–7].

Jet injectors are needle-free injection devices that can deliver liquids into the skin using a high-pressure stream. The recently introduced next-generation jet injectors are electronic pneumatic injection systems, which dispense fluids into the skin via accelerated, compressed gas. This minimally invasive and needle-free technique reduced occupational hazards, while providing drug delivery into the dermis leaving only a small entry point in the epidermis and a visible papule on the skin [8]. While the electronic pneumatic injection (EPI) device is more expensive and larger than spring-loaded jet injectors, clinical advantages of EPI include adjustable settings (pressure level and volume), ease of use in an outpatient clinic, minimal patient discomfort, and disposable nozzles and syringes to prevent contamination [9].

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Conflict of Interest Disclosures: All authors have completed and submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest and have disclosed the following: The EnerJet device was loaned to Merete Haedersdal by PerfAction as part of a research collaboration and the loan of *ex vivo* confocal microscopy was provided by Mavig.

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Optical imaging can be a valuable tool when the clinical utility of the various device parameters is explored. To investigate EPI parameters, skin-imitating media such as polyacrylamide gel and gelatin phantom have been used previously [10–12]. Moreover, EPI of ink followed by skin splitting and macroscopic evaluation, and histology imaging with hematoxylin and eosin (H&E) staining have also been used for optical imaging of EPI [10,11,13–15]. Although transparent media provide a clear view of injected fluid patterns, skin phantoms are currently lacking the technical refinement to mimic the biological complexity of human skin [16]. Macroscopic evaluation of skin tissue does display drug distribution but lacks detailed illustration of skin compartments and tissue alterations. Finally, cryosections can be used to inspect the cutaneous microstructure. The proneness to artifacts and inherent need for a multistep staining procedure to achieve sufficient image contrast, however, supports the use of novel optical imaging techniques to supplement traditional histology when assessing novel drug delivery techniques [17]. Fluorescence confocal microscopy can overcome these limitations by visualizing immediate EPI-induced fluorescent drug distribution without destructive tissue preparation or additional staining steps.

The *ex vivo* confocal microscopy (EVCN) uses a laser system to obtain high-resolution images of excised tissue [18]. The device used in the present study uses two lasers to generate both fluorescence (FCM) and reflectance confocal microscopy (RCM) images [19]. AO is most commonly used as contrast agent due to its high fluorescent yield, rapid staining and its well-established use in research [20]. EVCN is also being investigated to assess the margin control of basal cell carcinoma during micrographic Moh's surgery to shorten procedure time without sacrificing accuracy [21–28].

Visualization of immediate EPI-induced drug distribution at a microscopic level may provide a better understanding of its mechanism of action and clinical utility. Therefore, we aimed to visualize the immediate cutaneous drug distribution and changes in skin architecture using EVCN, and focus on the related clinical endpoint of EPI.

MATERIALS AND METHODS

Study Design

In this experimental *ex vivo* porcine study, immediate EPI-induced drug distribution was investigated with EVCN ($n = 22$ samples) and changes in skin architecture were investigated using EVCN and frozen histology ($n = 33$ samples). This study was conducted at the Department of Dermatology at the Bispebjerg Hospital in Copenhagen, Denmark.

Intervention

Flank skin samples from three Danish mixed-race Landrace/Yorkshire/Duroc pigs (3 months, 31–44 kg) stored at -80°C were thawed at room temperature for 30 minutes. Hair was trimmed using an electric razor. Skin samples were placed on plastic film surrounded by moist gauze for the

duration of the experiment. AO (MW: 301.8 Da, excitation wavelength = 500 nm, emission wavelength = 526 nm) (Mavig, Munich, Germany) was used as a contrast agent for EVCN [29]. The stock solution of 10 mg/ml was diluted in phosphate-buffered saline to a concentration of 0.06 mg/ml. Saline was used for control injections. EPI was performed with a needle-free electronic pneumatic injection device (EnerJet2.0; PerfAction Technologies Ltd., Rehovot, Israel) (Fig. 1C). The device has a fixed nozzle size of 200 μm . A volume of 50 μl was delivered per injection (device range; 50–150 μl) with pressure levels of 4 and 6 bar corresponding 50% and 100% pressures of the device (device range; 2–6 bar). Immediately after EPI, a caliper measured the diameter of the papular skin reaction by LB. Clinical photos were captured with a DSLR camera (EOS 750D; Canon) under standardized conditions.

Biopsy Processing and Evaluation of EVCN Images

Biopsy samples were bisected at the point of injection, placed on a glass slide, and flattened using a second slide and adhesive putty. The samples were mounted on an EVCN (Vivascope 2500; Mavig) to capture images of RCM (488 nm) and FCM (785 nm), respectively, at 50% and 15% intensity. EVCN provided automatic integration of RCM and FCM into pseudo-colored composite images. Resolution of the images was 1024 \times 1024 pixels.

Axial and lateral dispersion of AO in dermis were manually assessed by a nonblinded evaluator (LB) on FCM images. Axial distribution was defined as the perpendicular distance between the basal layer down to the deepest observed AO signal. Lateral distribution was defined as the maximum lateral expanse of AO signal, parallel to the epidermis.

Qualitative assessment of FCM, RCM, and composite images was performed by inspecting skin compartments, point of EPI entry, AO distribution pattern, and skin architecture disruption.

Evaluation of Frozen Histology

Samples for histological analyses were collected after EPI to assess changes in skin architecture. Cryosectioning, using a cryostat and freezing medium (Tissue-Tek[®] O.C.T.[™] Compound; Sakura Finetek Europe BV, Alphen, NL), was performed to harvest vertical sections at entry point of injection and of surrounding injection site with a slice thickness of 10 μm . Sections were stained with H&E and qualitatively evaluated (LB) under a light microscope with fourfold magnification.

Statistics

Mann–Whitney U tests compared papule diameter and immediate drug distribution parameters from EVCN images. Data were presented as median and interquartile ranges. An α level of $P < 0.05$ was considered statistically significant. Analyses were performed in SPSS version 25 (IBM Corporation; Armonk, NY).

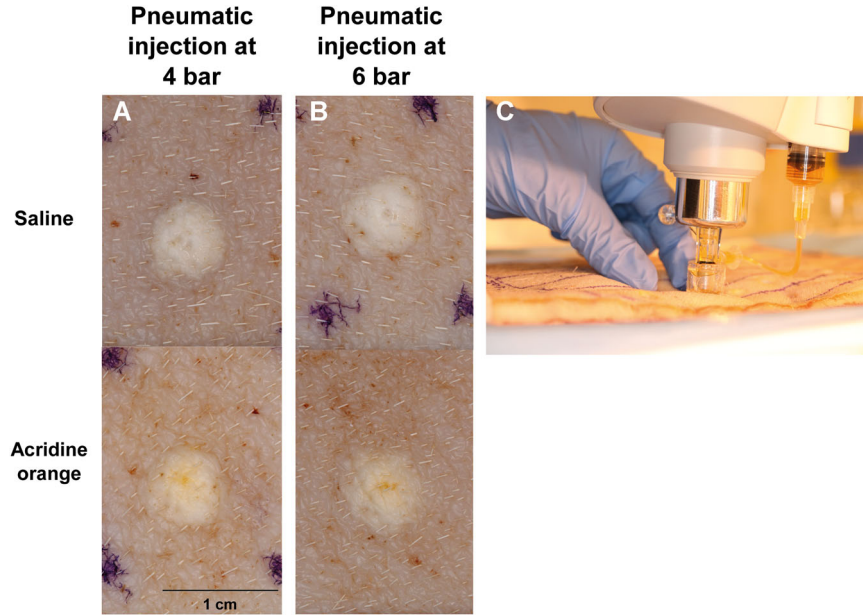


Fig. 1. Skin papule as clinical endpoint. A skin papule served as predefined, clinical endpoint after electronic pneumatic injection (50 μ l/injection) at 4 bar (A) and 6 bar (B). The pressure level did not have a significant effect on papule dimensions ($P=0.288$), with a diameter of 6.0 mm (6.0–7.0) at 4 bar and 7.0 mm at 6 bar (6.0–7.5 mm). The nozzle of the electronic pneumatic injection device is shown (C).

RESULTS

Papule as Clinical Endpoint

A distinct papule on the skin appeared after EPI ($n=55$). The pressure level was titrated up to 4 bar to consistently produce a papule after each injection. The pressure level did not appear to have a significant effect on papule dimensions ($n=33$; $P=0.288$), with a diameter of 6.0 mm (6.0–7.0) at 4 bar and 7.0 mm at 6 bar (6.0–7.5 mm; Fig. 1A and B).

Dermal Drug Distribution

EPI with 4 and 6 bar pressure levels resulted in a comparable dispersion of contrast agent reaching upper, mid, and deep dermis with an axial distribution of 2319 μ m (1500–2900 μ m; 4 bar) and 2399 μ m (2024–2758 μ m; 6 bar; $P=0.842$; Fig. 2; Table 1). Lateral dispersion for 4 and 6 bar pressure levels were also similar resulting in a lateral distribution of 5944 μ m (5173–6861 μ m; 4 bar) and 6187 μ m (5351–6544 μ m; 6 bar; $P=0.905$).

The most commonly observed distribution pattern resembled a hemispherical shape for both 4 and 6 bar pressure levels (Fig. 2A–C). However, variations in shape were observed between single injections resembling, for example, triangular deep-dermal or flat upper-dermal distribution patterns (Figs. 2D–F and 3). Overall, epithelium of hair follicles and epidermal entry point of injection showed high uptake of AO. The subcutaneous layer was reached sporadically.

Skin Architecture Disruption

Confocal microscopy images demonstrated disruption of normal skin structure after EPI with multiple vacuoles surrounding the dermal injection site and sub-epidermal clefting, independent of pressure levels (Fig. 3). The epidermis was torn apart at the entry point while architectural structure of the subcutaneous layer remained intact.

Tissue disruption after EPI was confirmed on histological sections displaying vacuoles, fragmented collagen, and tissue gaps in the dermis, to a similar degree at 4 and 6 bar pressure levels (Fig. 4). Disruption was most pronounced in sections closest to the point of entry. In contrast to confocal microscopy images, signs of epidermal trauma were not visible but sporadic disruption of subcutaneous tissue was seen.

DISCUSSION

We conducted an experimental *ex vivo* porcine study in which immediate EPI-assisted drug delivery was investigated using EVCM and histological sections. Our results demonstrate that a papule on normal skin is a reproducible clinical endpoint for EPI and indicates dermal drug delivery throughout all layers of the dermis, independent of pressure level settings. In laser and light treatments, clinical endpoints already have been shown to be helpful for safe and effective treatments; for example, immediate whitening of the skin after laser treatment using Q-switched lasers for tattoo removal [30]. In our study, we found that a skin papule has been consistently produced with a minimum pressure level of 4 bar. However, this

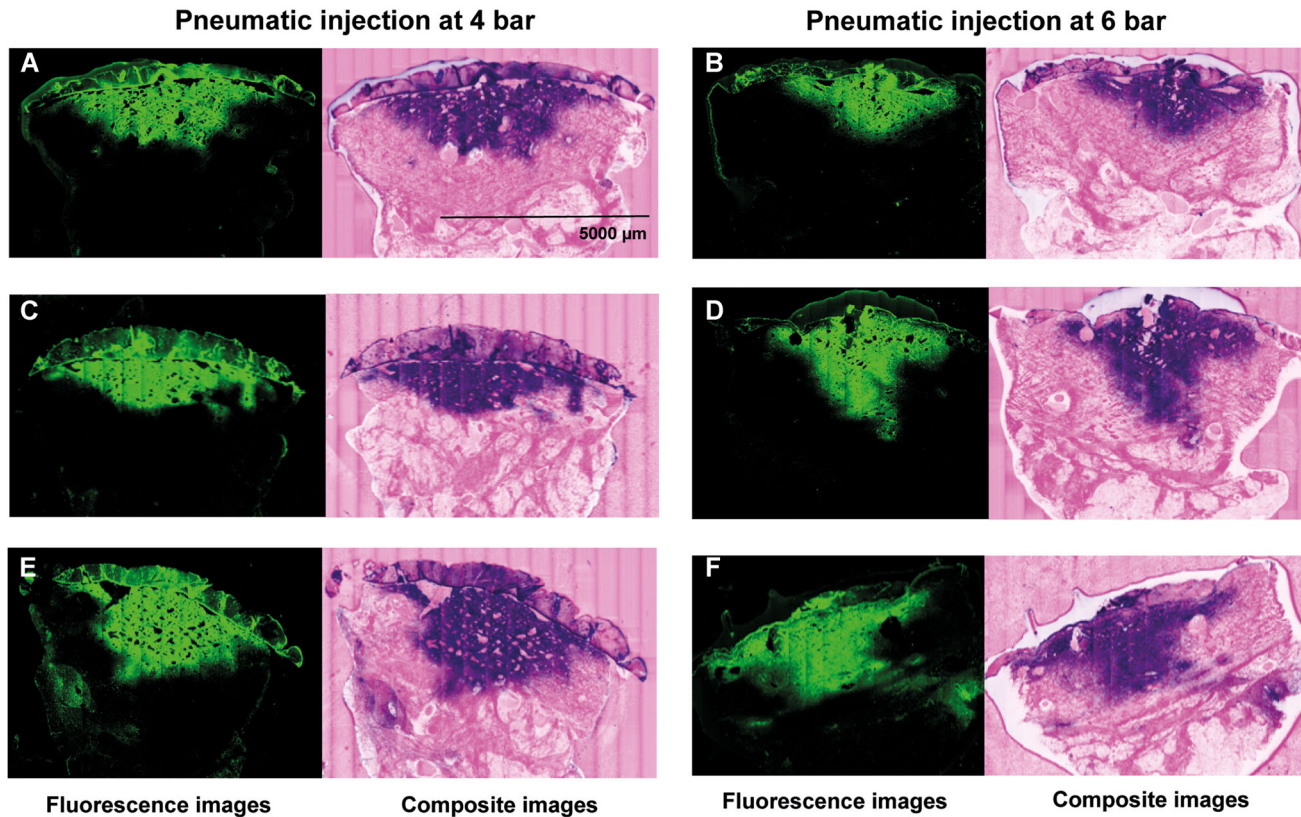


Fig. 2. Electronic pneumatic injection induces deep dermal drug delivery. Fluorescence and composite images were captured using an ex vivo confocal microscope. Images show comparable fluorescent drug distribution of acridine orange in deep dermis following electronic pneumatic injection at 4 and 6 bar pressure level (acridine orange solution is green on fluorescence images and purple on composite images). The most commonly observed distribution pattern was hemispherical (A, B, and C). Mild variations in shape have been observed for both pressure levels (D, E, and F). (A–F) display separate representative biopsies.

pressure level may not be directly translated into a clinical setting, as human dermis is generally thinner than porcine skin. Also, the pressure level required to induce a papule may vary when treating different anatomical locations or skin lesions with higher consistencies (e.g., hypertrophic scars or keloids).

In this study, we used EVCM as a high-resolution imaging technique. The EVCM had an acquisition time of

approximately 3–5 minutes, while preparation of cryosections requires 20–45 minutes per tissue sample [22]. Beside the benefit of minimal waiting time, tissue integrity remained intact and provided a realistic view on skin architecture and immediate drug distribution. Analyses of EVCM images showed that both 4 and 6 bar pressure levels generated deep dermal drug delivery. Spatial distribution of dermal drug delivery has also been investigated using a similar EPI device at 3.1, 3.9, and 4.6 bar pressure levels and 80 μ l injection volume by Erlandsson et al. [31], however, with ink as a contrast agent and assessment of drug distribution on histological sections only. In the study by Erlandsson et al., lateral dispersion was found to be pressure independent for individual injections, which is similar to our results. Notably, the lateral dispersion increased when stacking two consecutive injections at the same site, however, stacking of injections was not investigated in our study. For axial dispersion, a significant increase was observed when the highest pressure level of 4.6 bar was used for individual injections. This suggests that axial dispersion could be pressure dependent at lower pressure levels, while the dispersion is pressure independent between

TABLE 1. Spatial Distribution of Acridine Orange

Pressure level	Number of biopsies	Axial, median in μ m (IQR)	Lateral, median in μ m (IQR)
4 bar	10	2319 (1500–2900)	5944 (5173–6861)
6 bar	9	2399 (2024–2758)	6187 (5351–6544)

Spatial distribution of acridine orange in porcine dermis administered with electronic pneumatic injection, visualized by ex vivo confocal microscopy, defined as axial and lateral dermal drug distribution.

IQR, interquartile range.

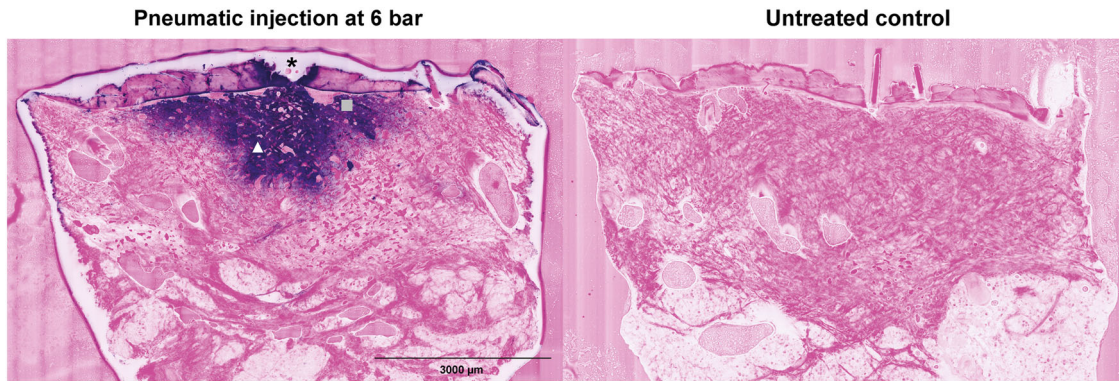


Fig. 3. Disrupted skin architecture by electronic pneumatic injection. Composite image showing skin architecture (pink) disrupted by electronic pneumatic injection with acridine orange (purple) at 6 bar pressure level (left) and an untreated biopsy (right) using ex vivo confocal microscopy. Tissue disruption was similar for 4 and 6 bar pressure level. *, epidermal trauma at entry point of injection; ■, clefting between stratum basale and dermis; ▲, vacuoles in the dermis.

4 and 6 bar according to our findings. Seok et al. [13] has found similar EPI-induced spatial distribution for 6 bar with an axial distribution of $2323\mu\text{m}$ compared with $2399\mu\text{m}$ found in this study. Although no difference in spatial distribution was observed between 4 and 6 bar pressure level, we did observe a variation in distribution

patterns, including hemispherical dispersion projecting from injection site, a deep-dermal distribution with a triangular appearance, and a flat upper-dermal distribution with a small penetration depth. This variation could possibly be explained by a difference in tissue consistencies within the skin sample and/or variation in

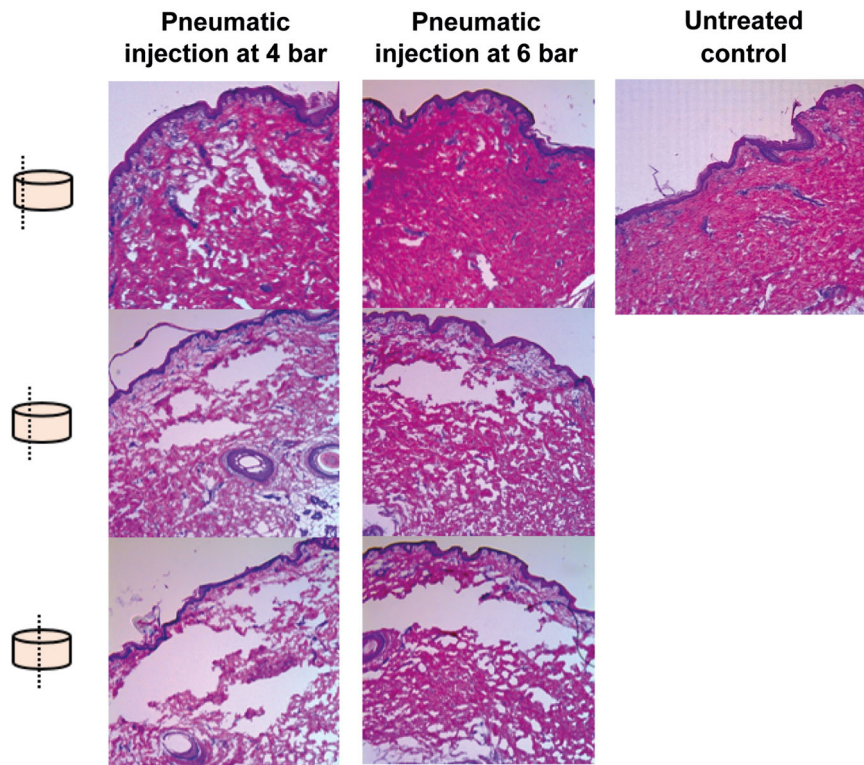


Fig. 4. Skin architecture disruption visualized on histology images. On frozen tissue sections, skin architecture disruption induced by electronic pneumatic injection was observed as vacuoles, fragmented collagen, and tissue gaps in the dermis. Three sections are shown of one representative biopsy for both 4 and 6 bar. The tissue disruptions are more prominent in close proximity to the point of entry. Untreated control with intact skin architecture is shown for comparison.

residue formation on top of the skin. Michinaka et al. [32] showed that the dispersion area of spring-driven jet injector relates inversely to the size of injected particles but proportionally to injection volume. Rohilla et al. [33] showed a significant effect on penetration depth, which could alter the distribution pattern, when comparing fluid viscosity *in vitro* using a spring-driven jet injector. The different working mechanisms of spring-driven jet injectors and EPI, however, preclude direct comparison of device parameters.

EPI-induced dermal skin structure disruption was visualized by EVCM images and histological sections, although with a slightly different presentation. The distinction could be explained by tissue alterations due to slow freezing, biopsy storage at -80°C , and tissue processing, which may have stretched the fragile tissue, exposing fragmented collagen and tissue gaps on histological sections [17]. Skin structure disruption of the dermis plays an important role in long-term effects of treatments focusing on skin remodeling. It activates the wound-healing process including the remodeling phase, which consists of collagen augmentation, crosslinking, and skin contraction for at least 6 months [12,15,34]. Previous studies have related EPI-induced neocollagenesis to a beneficial long-term effect of 4–6 months in rhytids [12,35], in contrast to the short-term effects seen after needle injection with, for example, hyaluronic acid and saline [35].

EPI's ability to provide deep dermal drug delivery may be particularly advantageous for dermatological conditions in which treatment specifically targets the dermis. In accordance, preliminary positive outcomes have been reported for keloids, acne scars, wrinkle treatment, palmar and axillar hyperhidrosis, local anesthetics, photo dynamic therapy for non-melanoma skin cancer, and nail psoriasis [36–41]. However, none of these studies focused on a clinical endpoint as in our study, which is important for guiding the treating physician in clinical practice.

Limitations in visualization of EPI-induced dermal drug delivery in this study includes the use of *ex vivo* pig skin to model *in vivo* human skin [42], uncertainty to what degree drugs used in clinical practice will behave similarly to AO, the lack of kinetic evaluation of cutaneous drug distribution, and the lacking assessment of a minimum threshold for AO detection with EVCM. Moreover, cryosectioning was used to assess tissue morphology with a moderate risk of artifacts, image assessment lacked a blinded second evaluator, and in order to standardize interventions, we only applied a fixed volume of 50 μl compared at 4 and 6 bar.

For future research we suggest to further investigate other factors that could impact the spatial dispersion of EPI, including different injection volumes, variation in residue formation on top of the skin, and stacking of injections.

CONCLUSION

This is the first study that relates a reproducible clinical endpoint to immediate EPI-assisted drug delivery and

tissue disruption using reflectance and fluorescence confocal microscopy images and histological sections. An EPI-induced skin papule indicates dermal drug delivery throughout all layers of the dermis, independent of pressure level settings.

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